	Consommation et Corporations Canada	Consumer and Corporate Affairs (	Canada 11)	2,005,059	
•	Bureau des brevets	Patent Office	(22)	1989/12/11	5,0
	Ottawa, Canada K1A 0C9		(43)	1990/06/12	073,
			(52)		8/90

# (19) (CA) APPLICATION FOR CANADIAN PATENT (12)

- (54) TNF Peptides
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- (30) (DE) P 38 41 759.6 1988/12/12
- (57) 8 Claims

Notice: The specification contained herein as filed

**Canadä** 

CCA 3254 (10-89) 41

O.Z.0050/40387

ABSTRACT OF THE DISCLOSURE: Peptides of the formula

X-A-Y,

where A, X and Y are defined in the description, and the preparation thereof are described. The novel peptides are suitable for controlling diseases.

O.Z.0050/40387

#### NOVEL THE PEPTIDES

The pr s nt invention relat s to novel p ptides derived from tumor necrosis factor (TNF), the preparation thereof and the use thereof as drugs.

Carswell et al. (Proc. Natl. Acad. Sci. USA 72 (1975) 3666) reported that the serum of endotoxin-treated animals which had previously been infected with the Calmette-Guerin strain of Mycobacteria (BCG) brought about hemorrhagic necrosis in various mouse tumors. This activity was ascribed to tumor necrosis factor. TNF also has a cytostatic or cytotoxic effect on a large number of transformed cell lines in vitro, whereas normal human and animal cell lines are unaffected (Lymphokine Reports Vol. 2, pp 235-275, Academic Press, New York, 1981). Recently, the biochemical characterization and the gene for human TNF have been described (Nature 312 (1984) 724, J. Biol. Chem. 260 (1985) 2345, Nucl. Acids Res. 13 (1985) 6361).

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It is possible to deduce from this data the following protein structure for mature human TNF:

ValargserserserärgfinrProseräspiyaProValalaHisValValalaAsnPro
GinalaGluGlyGlnieuGlnfrpieuAsnArgargalaAsnAlaleuleuAlaAsnGly
ValGluIeuArgaspasnGlnieuValValProserGluGlyIeuTyrieuIlefyrser
GinValleuPheilyeGlyGlnGlyCysProserThrHisValleuleufhrHisThrIle
SerärgIleAlaValSerTyrGlnfhriysValasnIeuleuSeralaIleIysSerPro
CysGlnärgGlufhrProGluGlyAlaGluAlaIysProTrpTyrGluProIleTyrIeu
GlyGlyValPheGlnieuGluIysGlyAspärgieuSeralaGluIleAsnArgProAsp
TyrIeuAspPheAlaGluSerGlyGlnValTyrPheGlyIleIleAlaleu

The TNF genes of cattle, rabbits and mice have also been described (Cold Spring Harbor Symp. Quant. Biol. <u>51</u> (1986) 597).

Besides its cytotoxic properti s, TNF is one of the main substances involved in inflammatory reactions (Pharmac. Res. <u>5</u> (1988) 129). Animal models have shown that TNF is involved in septic shock (Science <u>229</u> (1985) 869) and graft-versus-host disease (J. Exp. Med. <u>166</u> (1987) 1280).

We have now found that peptides with a considerably lower molecular weight have beneficial properties. The present invention relates to peptides of the formula

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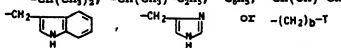
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X-A-Y I,

where

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- A is -Thr-Pro-Glu-Gly-Ala-, -Thr-Pro-Glu-Trp-Ala-,
  -Thr-Pro-Glu-Glu-Ala-,
  -Pro-Gly-Leu-Gln-Glu-Pro-, -Pro-Gly-Pro-Gly-Pro
  or -Pro-Gly-Leu-Gln-Gly-Pro-
- X is G-NH-CHM-CO-, G-NH-CHM-CO-W-, G-R-NH-CHM-CO- or G-R-NH-CHM-CO-W- and
- Y is -z, -NH-CHQ-CO-z, -V-NH-CHQ-CO-z, -NH-CHQ-CO-U-Z or -V-NH-CHQ-CO-U-Z,
- 20 where, in X and Y,
  - G is hydrogen or an amino-protective group,
  - Z is OH or NH2 or a carboxyl-protective group, or
  - G and Z together are also a covalent bond or -CO-(CH<sub>2</sub>)<sub>a</sub>-NH- where a is from 1 to 12,
- 25 R, U, V and W are peptide chains composed of 1-4 naturally occurring a-amino acids and
  - M and Q are hydrogens or one of the following  $-CH(CH_3)_2$ ,  $-CH(CH_3)-C_2H_5$ ,  $-C_9H_5$ ,  $-CH(OH)-CH_3$ ,



(with b being from 1 to 6 and T being hydrogen or OH, CH<sub>3</sub>O, CH<sub>3</sub>S, (CH<sub>3</sub>)<sub>2</sub>CH, C<sub>6</sub>H<sub>5</sub>, p-HO-C<sub>6</sub>H<sub>4</sub>, HS, H<sub>2</sub>N, HO-CO, H<sub>2</sub>N-CO or H<sub>2</sub>N-C(=NH)-NH) or

M and Q together are a  $-(CH_2)_c$ -S-S- $(CH_2)_d$ -,  $-(CH_2)_e$ -CO-NH- $(CH_2)_t$ - or  $-(CH_2)_e$ -NH-CO- $(CH_2)_t$ -NH-CO- $(CH_2)_t$ - bridge

(with c and d being from 1 to 4, e and f being from 1 to 6 and g being from 1 to 12), as well as the salts thereof with physiologically tolerated acids.

5 The peptides of the formula I are constructed of L-amino acids, but they can contain 1 or 2 D-amino acids. The side-chains of the trifunctional amino acids can carry protective groups or be unprotected.

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Particularly preferred physiologically tolerated acids are: hydrochloric acid, citric acid, tartaric acid, lactic acid, phosphoric acid, methanesulfonic acid, acetic acid, formic acid, maleic acid, fumaric acid, malic acid, succinic acid, malonic acid, sulfuric acid, L-glutamic acid, L-aspartic acid, pyruvic acid, mucic acid, benzoic acid, glucuronic acid, oxalic acid, ascorbic acid and acetylglycine.

The novel peptides can be open-chain (G = H, aminoprotective group; Z = OH,  $NH_2$ , carboxyl-protective group, M and Q not connected together) and, in particular, have a disulfide bridge (G = H, amino-protective group; Z = OH,  $NH_2$ , carboxyl-protective group;  $M + Q = -(CH_2)_0 - S-S-(CH_2)_0$ ) or a side-chain bridge (G = H, amino-protective group, Z = OH,  $NH_2$ , carboxyl-protective group,  $M + Q = -(CH_2)_0 - NH-CO-(CH_2)_1$  or  $-(CH_2)_0 - NH-CO-(CH_2)_1$  or  $-(CH_2)_0 - NH-CO-(CH_2)_2$  or  $-(CH_2)_0 - NH-CO-(CH_2)_3$  or be linked head-to-tail (G + Z = CO) covalent bond or  $-CO-(CH_2)_0 - NH-CO$ .

The novel compounds can be prepared by conventional methods of peptide chemistry.

Thus, the peptides can be constructed sequentially from amino acids or by linking together suitable smaller peptide fragments. In the sequential construction, the

p ptide chain is extended stepwis, by one amino acid each time, starting at the C terminus. In the case of coupling of fragments it is possible to link together fragments of different lengths, these in turn being obtainable by sequential construction from amino acids or coupling of other fragments. The cyclic peptides are obtained, after synthesis of the open-chain peptides, by a cyclization reaction carried out in high dilution.

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In the case both of sequential construction and of fragment coupling it is necessary for the building blocks to be linked by formation of an amide linkage. Enzymatic and chemical methods are suitable for this.

Chemical methods for forming amide linkages are dealt with in detail by Müller, Methoden der Organischen Chemie (Methods of Organic Chemistry) Vol. XV/2, pp 1-364, Thieme Verlag, Stuttgart, 1974; Stewart, Young, Solid Phase Peptide Synthesis, pp 31-34, 71-82, Pierce Chemical Company, Rockford, 1984; Bodanszky, Klausner, Ondetti, Peptide Synthesis, pp 85-128, John Wiley & Sons, New York, 1976 and other standard works of peptide chemistry. Particularly preferred are the azide method, the symmetrical and mixed anhydride method, active esters generated in situ or preformed and the formation of amide linkages using coupling reagents (activators), in particular dicyclohexylcarbodiimide (DCC), diisopropylcarbodiimide (DIC), 1-ethoxycarbonyl-2-ethoxy-1,2-dihydroquinoline (EEDQ), 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDCI), n-propanephosphonic anhydride N, N-bis(2-oxo-3-oxazolidinyl)amido-(PPA), phosphoryl chloride (BOP-Cl), diphenylphosphoryl azide (DPPA), Castro's reagent (BOP), O-benzotriazolyl-N,N,N',N'-tetra-methyluronium salts (HBTU), 2,5-diphenyl-2,3-dihydro-3-oxo-4-hydroxythiophene dioxide (Steglich's reagent; HOTDO) and 1,1'-carbonyldiimidazole (CDI). The c upling reagents can be employed alone or in combination with additives such as N,N'-dimethyl-4-aminopyridin (DMAP), N-hydroxybenzotriazole (HOBt), N-hydroxybenzotriazine (HOOBt), N-hydroxysuccinimide (HOSu) or 2-hydroxypyridine.

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Whereas it is normally possible to dispense with protective groups in enzymatic peptide synthesis, for chemical synthesis it is necessary for there to be reversible protection of the reactive functional groups which are not involved in the formation of the amide linkage on the two reactants. Three conventional protective group techniques are preferred for chemical peptide syntheses: the benzyloxycarbonyl (2), the t-butyloxycarbonyl (Boc) and the 9-fluorenylmethyloxycarbonyl (Fmoc) techniques. In each case the protective group on the a-amino group of the chain-extending building block is identified. The side-chain protective groups on the trifunctional amino acids are chosen so that they are not necessarily eliminated together with the a-amino protective group. A detailed review of amino acid protective groups is given by Müller, Methoden der Organischen Chemie Vol XV/1, pp 20-906, Thieme Verlag, Stuttgart, 1974.

The building blocks used to construct the peptide chain can be reacted in solution, in suspension or by a method similar to that described by Merrifield in J. Amer. Chem. Soc. 85 (1963) 2149. Particularly preferred methods are those in which peptides are constructed sequentially or by fragment coupling by use of the Z, Boc or Fmoc protective group technique, in which case the reaction takes place in solution, as well as those in which, similar to the Merrifield technique 3, one reactant is bound to an insoluble polymeric support (also called resin hereinafter). This typically entails the peptide being constructed sequentially on the polymeric support, by use of

the Boc or Fmoc protective group technique, with the growing peptide chain being covalently bonded at the C terminus to the insoluble resin particles (cf. Figures 1 and 2). This procedure allows reagents and byproducts to be removed by filtration, and thus recrystallization of intermediates is superfluous.

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The protected amino acids can be bonded to any suitable polymers which merely need to be insoluble in the solvents used and to have a stable physical form which allows easy filtration. The polymer must contain a functional group to which the first protected amino acid can be firmly linked by a covalent bond. A wide variety of polymers is suitable for this purpose, for example cellulose, polyvinyl alcohol, polymethacrylate, sulfonated polystyrene, chloromethylated copolymer of styrene and divinylbenzene (Merrifield resin), 4-methylbenzhydrylamine-resin (MBHA-resin), phenylacetamidomethyl-(Pam-resin), p-benzyloxybenzyl alcohol-resin, benzhydrylamine-resin (BHA-resin), 4-hydroxymethylbenzoyloxymethyl-resin, the resin used by Breipohl et al. (Tetrahedron Lett. 28 (1987) 565; from BACHEM), HYCRAM resin (from ORPEGEN) or SASRIN resin (from BACHEM).

Solvents suitable for peptide synthesis in solution are all those which are inert under the reaction conditions, in particular water, N,N-dimethylformamide (DMF), dimethyl sulfoxide (DMSO), acetonitrile, dichloromethane (DCM), 1,4-dioxane, tetrahydrofuran (THF), N-methyl-2-pyrrolidone (NMP) and mixtures of the said solvents. Peptide synthesis on polymeric supports can be carried out in all inert organic solvents which dissolve the amino acid derivatives used; however, solvents which also have resin-swelling properties are preferred, such as DMF, DCM, NMP, acetonitrile and DMSO, as well as mixtures of these solvents.

After the peptid has been synthesized it is cleaved off the polymeric support. The cleavage conditions for the various types of resins are disclosed in the literature. The cleavage reactions most commonly use acid and palladium catalysis, in particular cleavage in anhydrous liquid hydrogen fluoride, in anhydrous trifluoromethanesulfonic acid, in dilute or concentrated trifluoroacetic acid or palladium catalyzed cleavage in THF or THF-DCM mixtures in the presence of a weak base such as morpholine. The protective groups may, depending on the choice thereof, be retained or likewise cleaved off under the cleavage conditions. Partial deprotection of the peptide may also be worthwhile if the intention is to carry out certain derivatization reactions or a cyclization.

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Some of the novel peptides have good cytotoxic properties. Some others of the peptides have high affinity for the cellular TNF receptor without, however, having cytotoxic activity. They are therefore TNF antagonists. They compete with natural TNF for binding to the cellular TNF receptor and thus suppress the TNF effect. The novel peptides are valuable drugs which can be employed for treating neoplastic diseases and autoimmune diseases as well as for controlling and preventing infections, inflammations and transplant rejection reactions. Simple experiments can be used to elucidate the mode of action of the individual peptides. The cytotoxicity of the peptide is determined by incubating a TNP-sensitive cell line in the presence of the peptide. In a second experimental approach, the cell line is incubated with the relevant peptide in the presence of a lethal amount of It is possible in this way to detect the TNF-antagonistic effect. In addition, the affinity of the peptide for the cellular TNF receptor is determined in an in vitro binding experiment.

The following test systems wer used to charact rize the agonistic and antagonistic effects of the novel peptides:

- I. Cytotoxicity test on TNF-sensitive indicator cells,
- II. Cytotoxicity antagonism test on TNF-sensitive indicator cells,
- III. Competitive receptor-binding test on indicator cells expressing TNF receptor.

#### I. Cytotoxicity test

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The agonistic effects of the novel peptides are assessed on the basis of their cytotoxic effect on TNF-sensitive cells (e.g. L929, MCF-7, A204, U937). The test with L929 and MCF-7 was carried out as follows:

1. 100 μl of culture medium containing 3 to 5 x 10<sup>3</sup> freshly trypsinized, exponentially growing, L929 cells (mouse) or MCF-7 cells (human) were pipetted into the wells of a 96-well flat-bottom culture plate. The plate was incubated at 37°C overnight. The air in the incubator was saturated with water vapor and contained 5% CO<sub>2</sub> by volume.

The L929 culture medium contained 500 ml of 1x Earle's MEM (Boehringer Mannheim), 50 ml of heat-inactivated (56°C, 30 min) fetal calf serum (FCS), 50 ml of L-glutamine (200 mM), 5 ml of 100x non-essential amino acids, 3 ml of 1M HEPES buffer pH 7.2, and 50 ml of gentamicin (50 mg/ml).

The MCF-7 culture medium contained 500 ml of 1x Dulbecco's MEM (Boehringer Mannheim), 100 ml of heat-inactivated (56°C, 30 min) FCS, 5 ml of L-glutamine and 5 ml of 100x non-essential amino acids.

- 2. The next day 100 μl of the peptide solution to b tested were added to the cell cultures and subjected to serial 2-fold dilution. In addition, some cell controls (i. . c ll cultures not treated with peptide dilution) and some rhu-TNF controls (i.e. cell cultures treated with recombinant human TNF) were also made up. The culture plate was incubated at 37°C in an atmosphere of air saturated with water vapor and containing 5% CO<sub>2</sub> by volume for 48 h.
- 3. The percentage of surviving cells in the cultures treated with peptide dilution was determined by staining with crystal violet. For this purpose, the liquid was removed from the wells of the test plate by tapping it. 50 µl of crystal violet solution were pipetted into each well.

The composition of the crystal violet solution was as follows:

3.75 g of crystal violet

1.75 g of NaCl

161.5 ml of ethanol

43.2 ml of 37% formaldehyde

water ad 500 ml

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The crystal violet solution was left in the wells for 20 min and then likewise removed by tapping. The plates were then washed 5 times by immersion in water in order to remove dye not bound to the cells. The dye bound to the cells was extracted by adding 100  $\mu$ l of reagent solution (50% ethanol, 0.1% glacial acetic acid, 49.9% water) to each well.

- 4. The plat s were shak n for 5 min to obtain a solution of uniform color in each well. The surviving cells were determined by measuring the extinction at 540 nm of the colored solution in the individual wells.
- 5. Subsequently, by relating to the cell control, the 50% cytotoxicity value was defined, and the reciprocal of the sample dilution which resulted in 50% cytotoxicity was calculated as the cytotoxic activity of the test sample.

#### II. Cytotoxicity antagonism test

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The antagonistic effect of the peptides was assessed on the basis of their property of antagonizing the cytotoxic effect of rhu-TNF on TNF-sensitive cells (e.g. L929, MCF-7, A204, U937). The cytotoxicity antagonism test with L929 and MCF-7 cells was carried out as follows:

- 1. 100  $\mu$ l of culture medium containing 3 to 5 x 10<sup>3</sup> freshly trypsinized, exponentially growing, L929 cells (mouse) or MCF-7 cells (human) were pipetted into the wells of a 96-well flat-bottom culture plate. The plate was incubated at 37°C overnight. The air in the incubator was saturated with water vapor and contained 5% CO<sub>2</sub> by volume.
- The L929 culture medium contained 500 ml of lx Earle's MEM (Boehringer Mannheim), 50 ml of heat-inactivated (56°C, 30 min) FCS, 5 ml of L-gluta-mine (200 mM), 5 ml of 100x non-essential amino acids, 3 ml of 1M HEPES buffer pH 7.2, and 500 μl of gentamicin (50 mg/ml).

The MCF-7 culture medium contained 500 ml of 1x Dulbecco's MEM (Boehringer Mannheim), 100 ml of

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heat-inactivated (56°C, 30 min) FCS, 5 ml f L-glutamine (200 mM) and 5 ml of 100x non-essential amino acids.

- 2. The next day 100 μl of the peptide solution to be tested were added to the cell cultures and subjected to serial 2-fold dilution. Then, 100 μl of a rhu-TNF dilution in culture medium, which dilution had an 80-100% cytotoxic effect in the final concentration in the cell culture, were added to these cell cultures. In addition, some cell controls (i.e. cell cultures not treated with peptide solution or with rhu-TNF solution) and some rhu-TNF controls (= cell cultures treated only with rhu-TNF solution) were also made up. The culture plate was then incubated at 37°C in an atmosphere of air saturated with water vapor and containing 5% CO<sub>2</sub> by volume for 48 h.
- 3. The percentage of surviving cells in the cultures treated with substance solution was determined by staining with crystal violet. For this purpose, the liquid was removed from the wells of the test plate by tapping it. 50  $\mu$ l of crystal violet solution were pipetted into each well.

The crystal violet solution had the composition specified in I.3

The crystal violet solution was left in the wells for 20 min and then likewise removed by tapping. The plates were then washed 5 times by immersion in water in order to remove dye not bound to the cells. The dye bound to the cells was extracted by adding 100  $\mu$ l of reagent solution (50% ethanol, 0.1% glacial acetic acid, 49.9% water) to

each well.

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- 4. The plates were shaken for 5 min to obtain a solution of uniform color in ach well. The surviving cells were determined by measuring the extinction at 540 nm of the colored solution in the individual wells.
- 5. Subsequently, by relating to the cell control and the rhu-TNF control, the 50% antagonism value was defined, and the sample concentration which resulted in 50% antagonism of rhu-TNF cytotoxicity at the rhu-TNF concentration used was calculated as antagonistic activity of the sample tested.

# III. Competitive receptor-binding test

- Both the agonistic and antagonistic effects of peptides are conditional on the latter binding to the TNF receptor. This means that peptides with an agonistic or antagonistic effect compete with rhu-TNF for binding to the TNF receptor on TNF-sensitive indicator cells (e.g. U937). The competitive receptor-binding test was carried out as follows:
  - 1. 100 µl of medium containing various concentrations of the peptide to be tested and of rhu-TNF (= control) were pipetted into the reaction vessels. The medium comprised 500 ml of PBS (Boehringer Mannheim) containing 10 ml of heatinactivated (56°C, 30 min) FCS and 100 mg of sodium azide.
- 2. Subsequently, 100  $\mu$ l of medium containing 1 ng of <sup>125</sup>I-labeled rhu-TNF (Bolton lactoperoxidase

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method) were placed in the reaction vessels and mixed. The non-specific binding (NSB) was determined by mixing in the reaction vess ls the  $^{125}$ I-labeled rhu-TNF (1 ng of  $^{125}$ I-rhu-TNF in 100  $\mu$ l of medium) with a 200-fold excess of unlabeled rhu-TNF (200 ng of rhu-TNF in 100  $\mu$ l of medium).

- 3. Then 100  $\mu$ l of medium containing 2 x 10<sup>5</sup> U937 cells (human) were pipetted into the reaction vessels and mixed. The reaction vessels (test volume 300  $\mu$ l) were incubated at 0°C for 90 min. The reaction mixtures were remixed after 45 min.
- 4. After the incubation the cells were centrifuged at 1800 rpm and 4°C for 5 min, washed 3 times with medium and transferred quantitatively into counting vials, and the cell-bound radioactivity was determined in a Clini gamma counter 1272 (LKB Wallac).
- 5. After the measurements had been corrected for the non-specific binding, the 50% competition value was defined by relation to the overall binding, and the sample concentration which led to 50% competition of <sup>123</sup>I-rhu-TNF binding at the <sup>125</sup>I-rhu-TNF concentration used was calculated as the competitive activity of the sample tested.
- The Examples which follow are intended to explain the invention in more detail. The proteinogenous amino acids are abbreviated in the Examples using the conventional three-letter code. Other meanings are:

  Ac = acetic acid, Hcy = homocysteine, Orn = ornithine,

  Dap = 2,3-diaminopropionic acid.

## A. General procedur s

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I. The peptides claimed in claim 1 were synthesized using standard methods of solid-phase peptide synthesis in a completely automatic model 430A peptide synthesizer from APPLIED BIOSYSTEMS. The apparatus uses different synthesis cycles for the Boc and Fmoc protective group techniques.

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	a) Synthesis cycle for the Boc protects	ive group
	technique	
10	1. 30% trifluoroacetic acid in DCM	1 x 3 min
	2. 50% trifluoroacetic acid in DCM	1 x 17 min
	3. DCM washing	5 x 1 min
	4. 5% diisopropylethylamine in DCM	1 x 1 min
	5. 5% diisopropylethylamine in NMP	1 x 1 min
15	6. NMP washing	5 x 1 min
	7. Addition of preactivated protected amino	
	acid (activation by 1 equivalent of DCC	
	and 1 equivalent of HOBt in NMP/DCM);	
	peptide coupling (1st part)	1 x 30 min
20	8. Addition of DMSO to the reaction mixture	
	until it contains 20% DMSO by volume	
	9. Peptide coupling (2nd part)	1 x 16 min
	10. Addition of 3.8 equivalents of diiso-	
	propylethylamine to the reaction mixture	
25	11. Peptide coupling (3rd part)	1 x 7 min
	12. DCM washing	3 x 1 min
	13.If reaction is incomplete, repetition	
	of coupling (return to 5.)	
	14. 10% acetic anhydride, 5% diisopropyl-	
30	ethylamine in DCM	1 x 2 min
	15. 10% acetic anhydride in DCM	1 x 4 min
	16. DCM washing	4 x 1 min
	17 Return to 1.	

	b) Synt	thesis cycle for the Fmoc protective group	tech	mi	ďи	<b>a</b>
	1.	NMP washing	1	x	1	min
	2.	20% piperidine in NMP	1	x	4	min
	3.	20% piperidine in NMP	1	x	16	min
5	4.	NMP washing	5	×	1	min
	5.	Addition of preactivated protected amino				
		acid (activation by 1 equivalent of DCC				
		and 1 equivalent of HOBt in NMP/DCM);				
		peptide coupling	1	x	61	min
10	6.	NMP washing	3	x	1	min
	7.	If reaction is incomplete, repetition of				
		coupling (return to 5.)				
	8.	10% acetic anhydride in NMP	1	x	8	min
	9.	NMP washing	3	x	1	min
15	10.	Return to 2.				

Working up of peptide-resins obtained as in Ia The peptide-resin obtained as in Ia was dried under reduced pressure and transferred into a reaction vessel of a Teflon HF apparatus (from PENINSULA). Addition of a scavenger, preferably anisole (1 ml/g of resin), and of a thiol, in the case of tryptophancontaining peptides, to remove the indole formyl group, preferably ethanedithiol (0.5 ml/g of resin), was followed by condensation in of hydrogen fluoride (10 ml/g of resin) while cooling with liquid N2. The mixture was allowed to warm to 0°C, and was stirred at this temperature for 45 min. The hydrogen fluoride was then stripped off under reduced pressure and the residue was washed with ethyl acetate in order to remove remaining scavenger. The peptide was extracted with 30% strength acetic acid and filtered, and the filtrate was freeze-dried.

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To prepare peptide hydrazides, the peptide-resin (Pam- or Merrifield resin) was suspended in DMF

(15 ml/g of resin), hydrazine hydrate (20 equivalents) was added, and the mixture was stirred at room temperature for 2 days. T work up, the resin was filter d off and the filtrate was evaporated to dryn ss. The residue was crystalliz d from DMF/Et<sub>2</sub>O or MeOH/Et<sub>2</sub>O.

III. Working up of the peptide-resins obtained as in Ib

The peptide-resin obtained as in Ib was dried under reduced pressure and subsequently subjected to one of the following cleavage procedures, depending on the amino acid composition (Wade, Tregear, Howard Florey Fmoc-Workshop Manual, Melbourne 1985).

Peptide containing		Cleavage conditions				
15	Arg(Mtr)	Met	Trp	TFA	Scavenger	Reaction Time
	no	no.	no	95%	5% H <sub>2</sub> O	1.5 h
	yes	no	no	95%	5% thioanisole	≥ 3 h
20	no	yes	no	95%	5% ethyl methyl sulfide	1.5 h
	no	no	yes	95%	5% ethanedithiol/ anisole (1:3)	1.5 h
25	no	yes	yes	95%	5% ethanedithicl/ anisole/ethyl methy sulfide (1:3:1)	1.5 h
	yes	yes	yes	93%	7% ethanedithicl/ anisole/ethyl methy sulfide (1:3:3)	≥ 3 h
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The suspension of the peptide-resin in the suitable TFA mixture was stirred at room temperature for the

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stated tim and then th resin was filtered off and washed with TFA and with DCM. The filtrate and the washings w re extensively concentrated, and the peptide was precipitated by addition of diethyl ether. The mixture was cooled in an ice bath, and the precipitate was filtered off, taken up in 30% acetic acid and freeze-dried.

# IV. Purification and characterization of the peptides

Purification was by gel chromatography (SEPHADEX® G-10, G-15/10% HOAc; SEPHADEX® LH20/MeOH) and subsequent medium pressure chromatography (stationary phase: HD-SIL C-18, 20-45  $\mu$ , 100Å; mobile phase: gradient with A = 0.1% TFA/MeOH, B = 0.1% TFA/H<sub>2</sub>O).

The purity of the final products was determined by analytical HPLC (stationary phase:  $100 \times 2.1 \text{ mm}$  VYDAC C-18, 5  $\mu$ , 300 Å; mobile phase = CH<sub>3</sub>CN/H<sub>2</sub>O gradient buffered with 0.1% TFA, 40°C). Characterization was by means of amino acid analysis and fast atom bombardment mass spectrometry.

### 20 B. Specific procedures

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#### EXAMPLE 1

Ac-Ser-Pro-Thr-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-Pro-Trp-Tyr-NH2

0.98 g of Boc-Tyr (Br-Z) MBHA-resin (substitution 0.51 mmol/g), corresponding to a batch size of 0.5 mmol, was reacted as in AIa with 2 mmol each of

Boc-Trp(CHO)-OH	Boc-Gly-OH	Boc-Arg(Tos)-OH		
Boc-Pro-OH	Boc-Glu(OChx)-OH	Boc-Gln-OH		
Boc-Lys(C1-Z)-OH	Boc-Pro-OH	Boc-Thr(Bzl)-OH		

- 18 -

Boc-Ala-OH

B c-Thr(Bzl)-OH

Boc-Pro-OH

Boc-Glu(OChx)-OH

Boc-Glu(OCHx)-OH Boc-Ser(Bzl)-OH

Boc-Ala-OH

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After the synthesis was complete, the N terminus was acetylated (steps 1-6 and 14-16 as in AIa). The peptideresin was dried under reduced pressure; the yield was 2.2.g.

1.1 g of the resin obtained in this way were subjected to HF cleavage as in AII. The crude product (411 g) was purified by gel filtration (SEPHADEX® G-10) and medium pressure chromatography (cf. AIV; 50-65% A; 0.25% min<sup>-1</sup>). 242 mg of pure product were obtained.

#### EXAMPLE 2

H-Met-Val-Tyr-Pro-Gly-Leu-Gln-Glu-Pro-Trp-Leu-OH

0.47 g of Fmoc-Leu-p-alkoxybenzyl alcohol-resin (substit-15 ution 0.53 mmol/g), corresponding to a batch size of 0.25 mmol, was reacted as in AIb with 1 mmol each of

Fmoc-Trp-OH

Fmoc-Gly-OH

Fmoc-Pro-OH

Fmoc-Pro-OH

20 Fmoc-Glu(OtBu)-OH Fmoc-Tyr(tBu)-OH

Fmoc-Gln-OH

Fmoc-Val-OH

Fmoc-Leu-OH

Fmoc-Met-OH

After the synthesis was complete, the peptide-resin underwent N-terminal deprotection (steps 2-4 as in AIb). The resulting peptide-resin was dried under reduced pressure; the yield was 0.72 g.

The crude peptide (251 mg) obtained after TFA cleavage as in AIII was purified by gel filtration (SEPHADEX G - 10) and medium pressure chromatography (cf. AIV; 60-75% A; 0.25% min<sup>-1</sup>). 193 mg of pure product were obtained.

The following can be prepared in a similar manner to Examples 1 and 2:

- 3. H-Ser-Pro-Thr-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys- ::O-Trp-Tyr-OH
- 5 4. H-Ser-Pro-Tyr-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-Pro-Trp-Tyr-OH
  - 5. Ac-Ser-Pro-Tyr-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-Pro-Trp-Tyr-NH<sub>2</sub>
  - 6. Ac-Met-Val-Tyr-Pro-Gly-Leu-Gln-Glu-Pro-Trp-Leu-NH2

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#### EXAMPLE 7

Ac-Hcy-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-Pro-Hcy-NH<sub>2</sub>

0.98 g of Boc-Hcy(pMB)-MBHA-resin (substitution 0.51 mmol/g), corresponding to a batch size of 0.5 mmol, was reacted as in AIa with 2 mmol each of

	Boc-Pro-OH	Boc-Pro-OH	
	Boc-Lys(Cl-Z)-OH	Boc-Thr(Bzl)-OH	
20	Boc-Ala-OH	Boc-Glu(OChx)-OH	
	Boc-Glu-(OChx)-OH	Boc-Arg(Tos)-OH	
	Boc-Ala-OH	Boc-Gln-OH	
	Boc-Gly-OH	Boc-Hcy(pMB)-OH	
	Boc-Glu(OChx)-OH		

25 After the synthesis was complete, the N terminus was acetylated (steps 1-6 and 14-16 as in AIa). The peptideresin was dried under reduced pressure; the yield was 1.55 g.

0.78 g of the resin obtained in this way was subjected to HF cleavage as in AII. The freeze-dried crude product was taken up in 2 l of 0.1% strength acetic acid, and the pH was then adjusted to 8.4 with aqueous ammonia. Under an argon atmosphere, 0.01 N K<sub>3</sub>[Fe(CN)<sub>6</sub>] solution was slowly added dropwise until the yellowish-green color persisted for at least 15 min. The mixture was then stirred for 1 h and then acidified to pH 4.5 with glacial acetic acid, and 15 ml of an aqueous suspension of an anion exchanger (BIORAD 3 x 4A, chloride form) were added. After 30 min, the ion exchanger resin was filtered off, and the filtrate was concentrated to 100 ml in a rotary evaporator and subsequently freeze-dried.

All the solvents used had previously been saturated with nitrogen in order to prevent any oxidation of the free cysteine residues.

The crude product was purified by gel chromatography (SEPHADEX G-15) and medium pressure chromatography (cf. AIV; 40-60% A; 0.25% min<sup>-1</sup>). 58 mg of pure product were obtained.

The following can be prepared in a similar manner to Example 7 (Pam-resin was used to synthesize the peptide acids):

8. H-Cys-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-Pro-Cys-OH

9. Ac-Cys-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-

10. H-Hcy-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-

35 Pro-Cys-OH

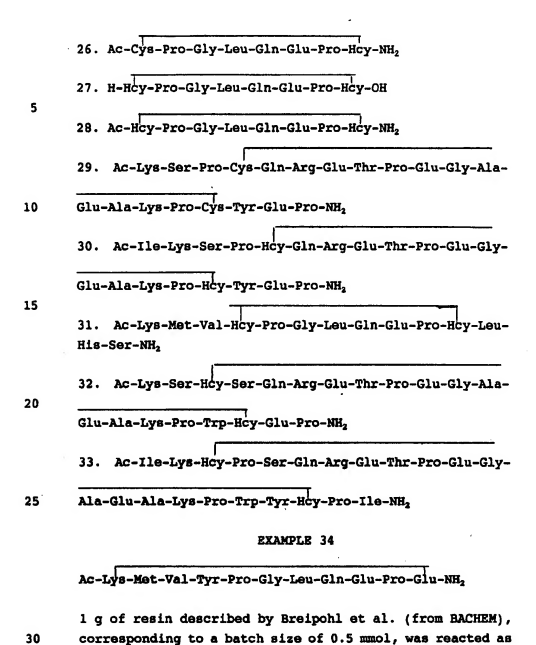
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	11. Ac-Hdy-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-
	Pro-Cys-NH <sub>2</sub>
5	12. H-Cys-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-
	Рго-нсу-Он
10	13. Ac-Cys-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-
	Pro-Hcy-NH <sub>2</sub>
	14. H-Hcy-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-
15	Pro-Hcy-OH
	15. Ac-Cys-Pro-Gly-Leu-Gln-Glu-Pro-Cys-NH <sub>2</sub>
20	16. H-Cys-Pro-Gly-Leu-Gln-Pro-Cys-OH
	17. H-Hcy-Pro-Gly-Leu-Gln-Glu-Pro-Cys-OH
25	18. Ac-Hcy-Pro-Gly-Leu-Gln-Glu-Pro-Cys-NH2
	19. H-Cys-Pro-Gly-Leu-Gln-Glu-Pro-Hcy-OH
	20. H-Cys-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Cys-NH2
30	21. Ac-Cys-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Cys-NH <sub>2</sub>
	22. H-Cys-Pro-Glu-Gly-Ala-Glu-Cys-NH2
a'e	23. Ac-Cys-Pro-Glu-Gly-Ala-Glu-Cys-NH <sub>2</sub>
35	24. H-Cys-Pro-Glu-Gly-Ala-Cys-NH <sub>2</sub>
	25. Ac-Cys-Pro-Glu-Gly-Ala-Cys-NH <sub>2</sub>



Fmoc-Glu(OtBu)-OH Fmoc-Leu-OH Fmoc-Val-OH Fmoc-Pro-OH Fmoc-Gly-OH Fmoc-Met-OH

in Alb with 2 mmol each of

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Fmoc-Glu(OBzl)-OH

Fmoc-Pro-OH

Fmoc-Lys(Boc)-OH

Fmoc-Gln-OH

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Fmoc-Tyr(tBu)-OH

After the synthesis was complete, the N terminus was acetylated (steps 2-4 and 8-9 as in AIb). The peptideresin was dried under reduced pressure; yield 1.86 g.

The crude product (615 mg) obtained after TFA cleavage as in AIII was dissolved in 500 ml of degassed DMF. 0.24 ml of NEt<sub>3</sub> and then, at -25°C, 0.24 ml of diphenylphosphoryl azide were added and the mixture was stirred at -25°C for 2 h. It was subsequently stored at -20°C for 2 days, at 4°C for 2 days and at room temperature for 2 days. It was then evaporated to dryness, and the crude peptide was purified by gel chromatography (SEPHADEX® LH 20). The isolated monomer (122 mg) was deprotected with HF as in AII and purified by medium pressure chromatography (cf. AIV, 45-60% A; 0.25% min<sup>-1</sup>). 83 mg of pure product were obtained.

#### EXAMPLE 35

20 <u>H-Ser-Ser-Gln-Lys-Met-Val-Tyr-Pro-Gly-Leu-Gln-Glu-Pro-Glu-Leu-His-Ser-OH</u>

2.63 g of Boc-Ser(Bzl)-Merrifield resin (substitution about 0.38 mmol/g), corresponding to a batch size of 1 mmol, were reacted as in AIb with 4 mmol each of

25	Boc-His(Z)-OH	Fmoc-Leu-OH	Fmoc-Met-OH
	Boc-Leu-OH	Fmoc-Gly-OH	Fmoc-Lys-(Boc)-OH
	Fmoc-Glu(OtBu)-OH	Pmoc-Pro-OH	Fmoc-Gln-OH
	Fmoc-Pro-OH	Fmoc-Tyr(tBu)-OH	Fmoc-Ser(Bzl)-OH
	Fmoc-Glu(OBzl)-OH	Fmoc-Val-OH	Fmoc-Ser(Bz1)-OH
30	Fmoc-Gln-OH		

Subsequently the t-butyl and Boc protective groups were cleaved off (steps 1-6 as in AIa). The cyclization on th resin took place in NMP with the addition of 1.77 g of BOP and 1.74 ml of disopropylethylamine (16 h). The peptide-resin underwent N-terminal deprotection (steps 2-4 as in AIb) and drying under reduced pressure. The yield was 3.51 g. The crude product obtained after HF cleavage as in AII was purified by gel filtration (SEPHADEX® G-25) and medium pressure chromatography twice (cf. AIV; 40 to 60% A; 0.25% min<sup>-1</sup>). 23 mg of pure product were obtained.

The following can be prepared in a similar manner to Examples 34 and 35:

- 36. Ac-Lys-Met-Val-Tyr-Pro-Gly-Leu-Gln-Glu-Pro-Glu-NH2
- 37. Ac-Lys-Met-Val-Tyr-Pro-Gly-Leu-Gln-Glu-Pro-Glu-OH
- 38. Ac-Lys-Ser-Pro-Thr-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-
- 20 Glu-Ala-Lys-Pro-Trp-Tyr-Asp-NH<sub>2</sub>

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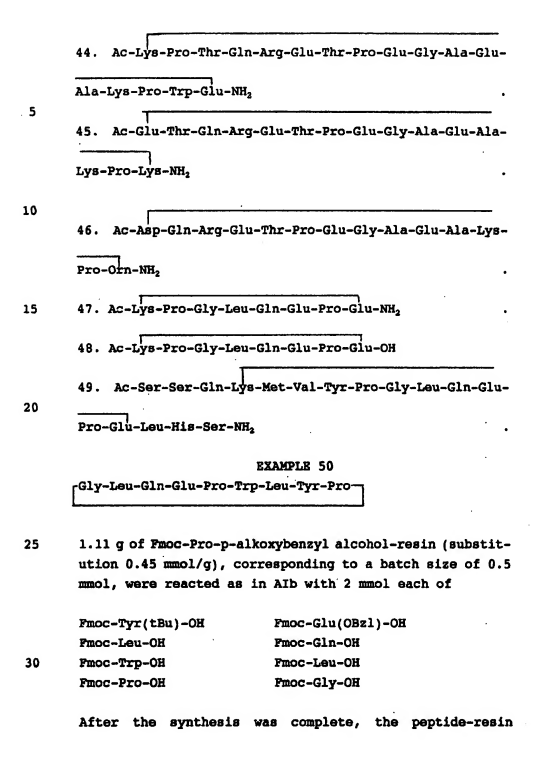
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- 39. Ac-Lys-Thr-Pro-Glu-Gly-Ala-Asp-NH<sub>2</sub>
- 40. H-Lys-Thr-Pro-Glu-Gly-Ala-Asp-OH
- 41. Ac-Glu-Thr-Pro-Glu-Gly-Ala-Lys-OH
- 42. Ac-Lys-Ser-Pro-Tyr-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-
- 30 Glu-Ala-Lys-Pro-Trp-Tyr-Glu-NH<sub>2</sub>
  - 43. Ac-Orn-Ser-Pro-Tyr-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-

Glu-Ala-Lys-Pro-Trp-Tyr-Asp-NH2



underwent N-terminal deprotection (steps 2-4 as in Alb), and subsequent drying und r reduced pressure. The yi ld was 1.55 g.

The crude peptide (486 mg) obtained after TFA cleavage as in AIII was dissolved in 500 ml of degassed DMF. 210 mg of NaHCO<sub>3</sub> and then, at - 25°C, 0.24 ml of diphenylphosphoryl azide were added, and the mixture was stirred at - 25°C for 2 hours and at room temperature for 2 days. It was then evaporated to dryness, and the crude peptide was purified by gel chromatography (SEPHADEX® LH 20). The isolated monomer (73 mg) was deprotected with HF as in AII and purified by medium pressure chromatography (cf. AIV; 60-70% A; 0.25% min<sup>-1</sup>). 32 mg of pure product were obtained.

- The following can be prepared in a similar manner to Example 50:
  - 51. <u>Tvr-Gln-Arg-Glu-Thr-Pro-Glu-Gly-Ala-Glu-Ala-Lys-Pro</u>

#### -Trp-

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- 52. <u>Lvs-Met-Val-Tvr-Pro-Gly-Leu-Gln-Glu-Pro-Glu-</u>
- 20 53. Tyr-Pro-Gly-Leu-Gln-Glu-Pro-Trp-

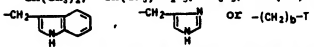
We claim: -

A peptide of the formula I,

#### X-A-Y I,

#### where

- A is -Thr-Pro-Glu-Gly-Ala-, -Thr-Pro-Glu-Trp-Ala-, -Thr-Pro-Glu-Glu-Ala-, -Pro-Gly-Leu-Gln-Glu-Pro-, -Pro-Gly-Pro-Gly-Pro- or -Pro-Gly-Leu-Gln-Gly-Pro-
- X is G-NH-CHM-CO-, G-NH-CHM-CO-W-, G-R-NH-CHM-CO- or G-R-NH-CHM-CO-W- and
- Y is -z, -NH-CHQ-CO-z, -V-NH-CHQ-CO-z, -NH-CHQ-CO-U-z or -V-NH-CHQ-CO-U-z, where, in X and Y,
- G is hydrogen or an amino-protective group,
- Z is OH or NH<sub>2</sub> or a carboxyl-protective group, or
- G and Z together are also a covalent bond or -CO-(CH<sub>2</sub>)<sub>a</sub>-NH- where a is from 1 to 12,
- R, U, V and W are peptide chains composed of 1-4 naturally occurring  $\alpha$ -amino acids and



(with b being from 1 to 6 and T being hydrogen or OH,  $CH_3O$ ,  $CH_3S$ ,  $(CH_3)_2CH$ ,  $C_6H_5$ ,  $p-HO-C_6H_4$ , HS,  $H_2N$ , HO-CO,  $H_2N-CO$  or  $H_2N-C(=NH)-NH$ ) or

M and Q together are a  $-(CH_2)_0$ -S-S- $(CH_2)_d$ -,  $-(CH_2)_s$ -CO-NH- $(CH_2)_s$ - or  $-(CH_2)_s$ -NH-CO- $(CH_2)_s$ -NH-CO- $(CH_2)_s$ - bridge (with c and d being from 1 to 4, e and f being from 1 to 6 and g being from 1 to 12),

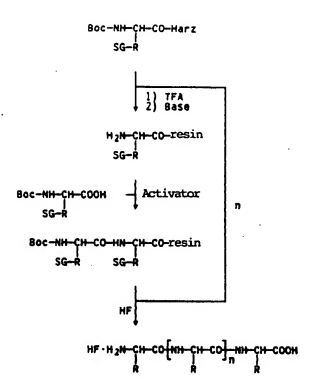
as well as the salts thereof with physiologically tolerated acids.

A peptide as claimed in claim 1, where G is hydrogen

or an amino-protective group and Z is hydroxyl or amino or a carboxyl-protective group, and M and Q ar not connected together.

- 3. A peptide as claimed in claim 1, where G is hydrogen or an amino-protective group and Z is hydroxyl or amino or a carboxyl-protective group, and M and Q together are a  $-(CH_2)_{\circ}-S-S-(CH_2)_{d}$  bridge.
- 4. A peptide as claimed in claim 1, where G is hydrogen or an amino-protective group and Z is hydroxyl or amino or a carboxyl-protective group, and M and Q together are -(CH<sub>2</sub>)<sub>e</sub>-NH-CO-(CH<sub>2</sub>)<sub>f</sub>- or -(CH<sub>2</sub>)<sub>e</sub>-NH-CO-(CH<sub>2</sub>)<sub>e</sub>-NH-CO-(CH<sub>2</sub>)<sub>f</sub>.
- A peptide as claimed in claim 1, where G + 2 together are a covalent bond or -CO-(CH<sub>2</sub>)<sub>a</sub>-NH-.
- 6. A peptide as claimed in claim 1 to 5 for use for controlling diseases.
- 7. The use of a peptide as claimed in claims 1 to 5 for controlling neoplastic diseases and autoimmune diseases as well as for controlling and preventing infections, inflammations and transplant rejection reactions.
- 8. A process for the preparation of a peptide as claimed in claims 1 to 5, which comprises preparation thereof using conventional methods of peptide chemistry.

# Fig. 1: The Boc prot ctive group technique on a polymeric support



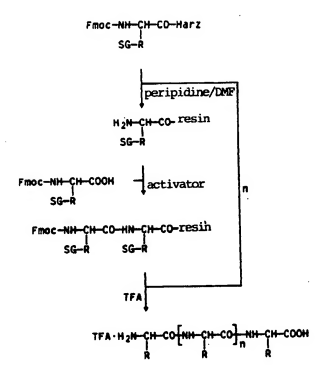
Boc = t-butyloxycarbonyl protective group

SG = side-chain protective group

R = amino acid side-chain

Mr. Mel

Fig. 2: Th Fmoc protective group technique on a polymeric support



Fmoc = 9-fluorenylmethoxycarbonyl protective group

SG = side-chain protective group

R = amino acid side-chain,

Patent Agents.